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                 Web Page for STN Seminar Schedule - N. America
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NEWS 3 JUL 02 SCISEARCH enhanced with complete author names
NEWS 4 JUL 02 CHEMCATS accession numbers revised
      5 JUL 02 CA/CAplus enhanced with utility model patents from China
NEWS
NEWS 6 JUL 16 CAplus enhanced with French and German abstracts
      7 JUL 18
                 CA/CAplus patent coverage enhanced
NEWS
NEWS 8 JUL 26 USPATFULL/USPAT2 enhanced with IPC reclassification
         JUL 30
                USGENE now available on STN
NEWS 9
NEWS 10 AUG 06 CAS REGISTRY enhanced with new experimental property tags
NEWS 11 AUG 06 FSTA enhanced with new thesaurus edition
NEWS 12 AUG 13 CA/CAplus enhanced with additional kind codes for granted
                 patents
NEWS 13 AUG 20 CA/CAplus enhanced with CAS indexing in pre-1907 records
NEWS 14 AUG 27 Full-text patent databases enhanced with predefined
                 patent family display formats from INPADOCDB
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                 spectral property data
         SEP 07
                 STN AnaVist, Version 2.0, now available with Derwent
NEWS 17
                 World Patents Index
NEWS 18
         SEP 13
                 FORIS renamed to SOFIS
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                 INPADOCDB enhanced with monthly SDI frequency
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                 CA/CAplus enhanced with printed CA page images from
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         SEP 17
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                 patents
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                 EMBASE, EMBAL, and LEMBASE reloaded with enhancements
         OCT 02
                 CA/CAplus enhanced with pre-1907 records from Chemisches
 NEWS 23
                 Zentralblatt
 NEWS 24 OCT 19
                 BEILSTEIN updated with new compounds
              19 SEPTEMBER 2007: CURRENT WINDOWS VERSION IS V8.2,
 NEWS EXPRESS
              CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
              AND CURRENT DISCOVER FILE IS DATED 19 SEPTEMBER 2007.
```

11/564,974

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SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

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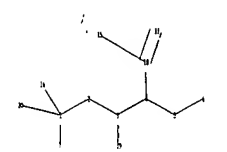
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REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

=> Uploading C:\Program Files\Stnexp\Queries\10564974.str



G1:H,Cy,Ak

G2:H,Ak

Match level:

1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:Atom 7:CLASS 10:CLASS 11:CLASS 15:CLASS 16:CLASS 19:CLASS 20:CLASS

L1 STRUCTURE UPLOADED

=> s 11 .

SAMPLE SEARCH INITIATED 13:02:15 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 580 TO ITERATE

100.0% PROCESSED 580 ITERATIONS

1 ANSWERS

28 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 10156 TO 13044

PROJECTED ANSWERS: 1 TO 80

L2 1 SEA SSS SAM L1

=> s 11 ful

FULL SEARCH INITIATED 13:02:24 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 12233 TO ITERATE

100.0% PROCESSED 12233 ITERATIONS

SEARCH TIME: 00.00.01

L3 28 SEA SSS FUL L1

=> file caplus

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FULL ESTIMATED COST 172.10 172.31

FILE 'CAPLUS' ENTERED AT 13:02:34 ON 09 NOV 2007 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2007 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 9 Nov 2007 VOL 147 ISS 21 FILE LAST UPDATED: 8 Nov 2007 (20071108/ED)

Effective October 17, 2005, revised CAS Information Use Policies apply. They are available for your review at:

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http://www.cas.org/infopolicy.html
=> s 13
L4
             4 L3
=> d l4 ibib abs hitstr hitind 1-4
                    CAPLUS COPYRIGHT 2007 ACS on STN
    ANSWER 1 OF 4
L4
ACCESSION NUMBER:
                         2005:411090
                                      CAPLUS
DOCUMENT NUMBER:
                         143:97618
TITLE:
                         HIV protease inhibitors: synthesis and activity of
                         N-aryl-N'-hydroxyalkyl hydrazide pseudopeptides
AUTHOR(S):
                         Marastoni, M.; Baldisserotto, A.; Trapella, C.;
                         McDonald, J.; Bortolotti, F.; Tomatis, R.
CORPORATE SOURCE:
                         Department of Pharmaceutical Sciences and
                         Biotechnology Center, University of Ferrara,
Ferrara,
                         I-44100, Italy
                         European Journal of Medicinal Chemistry (2005),
SOURCE:
40(5),
                         445-451
                         CODEN: EJMCA5; ISSN: 0223-5234
PUBLISHER:
                         Elsevier Ltd.
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         English
OTHER SOURCE(S):
                         CASREACT 143:97618
     We describe the synthesis by solution peptide coupling, N-acylation,
AB
and
     N-alkylation and activities of a series of pseudopeptides containing an
     N-aryl-N'-hydroxyalkyl hydrazide core structure to inhibit human
     immunodeficiency virus protease and viral replication. Of the series,
     compound Hmb-Leu-N(CH2Ph)-N(CH2-CH-OH)-rPro-Boc (Hmb = 3-hydroxy-2-
     methylbenzoyl, rPro = proline residue in the opposite direction, Boc =
     tert-butoxycarbonyl) displayed the greatest inhibitory potency (IC50 <
1
     \muM) and exhibited enzymic resistance and stability in vitro.
     856667-69-1P 856667-73-7P 856667-77-1P
IT
     RL: PAC (Pharmacological activity); SPN (Synthetic preparation); BIOL
     (Biological study); PREP (Preparation)
        (synthesis and activity of arylhydroxyalkyl hydrazide
pseudopeptides as
        HIV protease and viral replication inhibitors)
RN
     856667-69-1 CAPLUS
     1,2-Pyrrolidinedicarboxylic acid, 1-(1,1-dimethylethyl) ester,
CN
     2-[1-(2-hydroxyethyl)-2-[(2S)-2-[(3-hydroxy-2-methylbenzoyl)amino]-4-
     (methylthio) -1-oxobutyl] -2-(phenylmethyl) hydrazide], (2S) - (9CI)
                                                                       (CA
     INDEX NAME)
```

RN 856667-73-7 CAPLUS

CN 1,2-Pyrrolidinedicarboxylic acid, 1-(1,1-dimethylethyl) ester, 2-[1-(2-hydroxyethyl)-2-[(2S)-2-[(3-hydroxy-2-methylbenzoyl)amino]-4-methyl-1-oxopentyl]-2-(phenylmethyl)hydrazide], (2S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 856667-77-1 CAPLUS

CN 1,2-Pyrrolidinedicarboxylic acid, 1-(1,1-dimethylethyl) ester,
2-[1-(2-hydroxyethyl)-2-[(2S,3R)-3-hydroxy-2-[(3-hydroxy-2-methylbenzoyl)amino]-1-oxobutyl]-2-(phenylmethyl)hydrazide], (2S)-(9CI)

(CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

```
CC
    34-3 (Amino Acids, Peptides, and Proteins)
     Section cross-reference(s): 1, 7
     856667-69-1P 856667-71-5P 856667-72-6P 856667-73-7P
     856667-76-0P 856667-77-1P
     RL: PAC (Pharmacological activity); SPN (Synthetic preparation); BIOL
     (Biological study); PREP (Preparation)
        (synthesis and activity of arylhydroxyalkyl hydrazide
pseudopeptides as
        HIV protease and viral replication inhibitors)
REFERENCE COUNT:
                               THERE ARE 29 CITED REFERENCES AVAILABLE FOR
                         29
THIS
                               RECORD. ALL CITATIONS AVAILABLE IN THE RE
FORMAT
    ANSWER 2 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN
L4
ACCESSION NUMBER:
                         2005:120872 CAPLUS
DOCUMENT NUMBER:
                         142:219049
TITLE:
                         Preparation of benzoic acid derivatives having
                         hydrazide moiety as prostaglandin receptors
modulators
INVENTOR(S):
                         Araldi, Gian Luca; Liao, Yihua; Brugger, Nadia
PATENT ASSIGNEE(S):
                         Applied Research Systems Ars Holding N.V., Neth.
SOURCE:
                         PCT Int. Appl., 73 pp.
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English -
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                            APPLICATION NO.
     PATENT NO.
                         KIND DATE
                                                                   DATE
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•																		
		2005012232							WO 2004-EP51531						20040716			
	WO	2005012232			A3	A3 20050331			_									
		W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
			GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,
			LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,
			NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
			TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
		RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,
			AZ,	BY,	KG,	KZ,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,
			EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,	IT,	LU,	MC,	NL,	PL,	PT,	RO,	SE,
			SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,
			SN,	TD,	TG													
	AU								AU 2004-261397 CA 2004-2529123						20040716			
	CA														20040716			
	ΕP				A2	A2 20060510				EP 2004-785978					20040716			
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		•	IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	HU,	PL,	
SK,	HR																	
	JP	JP 2006528154			${f T}$	20061214				JP 2006-520831					20040716			

NO 2006000739

(CA INDEX NAME)

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US 2007185191
                                20070809
                                            US 2006-564974
                          A1
                                                                    20060711
PRIORITY APPLN. INFO.:
                                            US 2003-488614P
                                                                    20030718
                                                                    20040716
                                            WO 2004-EP51531
OTHER SOURCE(S):
                        CASREACT 142:219049; MARPAT 142:219049
GI
* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *
     Title compds. I [A = cycloalkyl, heterocycloalkyl, heteroaryl, etc.; B
AB
     alkylene, alkenylene; alkynylene; R1 = H, alkyl, alkenyl, etc.; R2, R3
     H, alkyl, alkenyl, etc.; R4 = H, alkyl; R5 = H, alkyl, alkenyl, etc.;
n =
     1-6] and their pharmaceutically acceptable salts were prepared For
example,
     Michael addition of the hydrazino compound II to compound III followed
by reduction
     with NaBH4, hydrolysis using NaOH afforded compound IV.
prostaglandin
     EP2 binding assays, the Ki value of compound IV was 4.21 \muM.
                                                                    Compds.
Ι
     are claimed useful for the treatment of asthma, hypertension, etc.
     841234-41-1P 841234-42-2P 841234-43-3P
     841234-44-4P 841234-45-5P 841234-46-6P
     841234-47-7P 841234-49-9P 841234-50-2P
     841234-51-3P 841234-52-4P 841234-53-5P
     841234-54-6P 841234-55-7P 841234-56-8P
     841234-57-9P 841234-58-0P 841234-59-1P
     841234-60-4P 841234-61-5P
     RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
     (Uses)
        (preparation of benzoic acid derivs. having hydrazide moiety as
        prostaglandin receptors modulators for treatment of asthma,
        hypertension, etc.)
     841234-41-1 CAPLUS
RN
     Benzoic acid, 4-[2-[1-acetyl-2-(3-hydroxybutyl)hydrazino]ethyl]- (9CI)
CN
```

20060331

A

NO 2006-739

20060215

$$\begin{array}{c} \text{NH-CH}_2\text{-CH}_2\text{-CH-Me} \\ \\ \text{CH}_2\text{-CH}_2\text{-N-Ac} \\ \\ \text{HO}_2\text{C} \end{array}$$

RN 841234-42-2 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-(3-hydroxyoctyl)hydrazino]ethyl]- (9CI) (CA INDEX NAME)

NH- CH₂- CH₂- CH- (CH₂)
$$_4$$
- Me CH₂- CH₂- N- Ac HO₂C

RN 841234-43-3 CAPLUS

CN Benzoic acid,

4-[2-[1-acetyl-2-(3-hydroxyoctyl)-2-methylhydrazino]ethyl]-(9CI) (CA INDEX NAME)

RN 841234-44-4 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-[3-hydroxy-4-[3-(trifluoromethyl)phenyl]butyl]hydrazino]ethyl]- (9CI) (CA INDEX NAME)

RN 841234-45-5 CAPLUS

CN Benzoic acid,

4-[2-[2-[3-hydroxy-4-[3-(trifluoromethyl)phenyl]butyl]-1-(1-oxopropyl)hydrazino]ethyl]- (9CI) (CA INDEX NAME)

RN 841234-46-6 CAPLUS

CN Benzoic acid,

4-[2-[2-[3-hydroxy-4-[3-(trifluoromethyl)phenyl]butyl]-1-(2-methyl-1-oxopropyl)hydrazino]ethyl]- (9CI) (CA INDEX NAME)

RN 841234-47-7 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-[3-hydroxy-4-[3-(phenylethynyl)phenyl]butyl]hydrazino]ethyl]- (9CI) (CA INDEX NAME)

RN 841234-49-9 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-(4-[1,1'-biphenyl]-3-yl-3-hydroxybutyl)hydrazino]ethyl]- (9CI) (CA INDEX NAME)

Ph
$$CH_2-CH-CH_2-CH_2-NH-N-CH_2-CH_2$$

RN 841234-50-2 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-[4-(3-bromophenyl)-3-hydroxybutyl]hydrazino]ethyl]- (9CI) (CA INDEX NAME)

RN 841234-51-3 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-[3-hydroxy-4-(3-iodophenyl)butyl]hydrazino]ethyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Ac} & \text{OH} \\ \text{CH}_2\text{--}\text{CH}_2\text{--}\text{NH--}\text{CH}_2\text{--}\text{CH}_2\text{--}\text{CH}_2\text{--}\text{CH}_2\\ \\ \text{HO}_2\text{C} & \end{array}$$

RN 841234-52-4 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-[4-(3-chlorophenyl)-3-hydroxybutyl]hydrazino]ethyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Ac} & \text{OH} \\ \text{CH}_2\text{--}\text{CH}_2\text{--}\text{NH}\text{--}\text{CH}_2\text{--}\text{CH}\text{--}\text{CH}_2\text{--}\text{CH}\text{--}\text{CH}_2\\ \text{HO}_2\text{C} & \\ \end{array}$$

RN 841234-53-5 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-(3-cyclohexyl-3-hydroxypropyl)hydrazino]ethyl]- (9CI) (CA INDEX NAME)

RN 841234-54-6 CAPLUS

CN Benzoic acid,

4-[2-[1-acetyl-2-(3-hydroxy-4-phenylbutyl)hydrazino]ethyl](9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{NH-CH}_2\text{--CH}_2\text{--CH-CH}_2\text{--Ph} \\ \\ \text{CH}_2\text{--CH}_2\text{--N-Ac} \\ \\ \text{HO}_2\text{C} \end{array}$$

RN 841234-55-7 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-[4-(4-chlorophenyl)-3-hydroxybutyl]hydrazino]ethyl]- (9CI) (CA INDEX NAME)

RN 841234-56-8 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-[4-(4-fluorophenyl)-3-hydroxybutyl]hydrazino]ethyl]- (9CI) (CA INDEX NAME)

RN 841234-57-9 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-[4-(3-ethynylphenyl)-3-hydroxybutyl]hydrazino]ethyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Ac} & \text{OH} \\ \text{CH}_2\text{--}\text{CH}_2\text{--}\text{N--}\text{NH--}\text{CH}_2\text{--}\text{CH--}\text{CH}_2 \\ \end{array}$$

RN 841234-58-0 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-[4-(3-fluorophenyl)-3-hydroxybutyl]hydrazino]ethyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Ac} & \text{OH} \\ \text{CH}_2\text{--}\text{CH}_2\text{--}\text{NH--}\text{CH}_2\text{--}\text{CH}_2\text{--}\text{CH}_2\text{--}\text{CH}_2\\ \text{HO}_2\text{C} \end{array}$$

RN 841234-59-1 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-[3-hydroxy-4-[4- (phenylethynyl)phenyl]butyl]hydrazino]ethyl]- (9CI) (CA INDEX NAME)

HO2C
$$AC$$
 OH $CH_2-CH_2-N-NH-CH_2-CH_2-CH-CH_2$

RN 841234-60-4 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-[3-hydroxy-4-(2-thienyl)butyl]hydrazino]ethyl]- (9CI) (CA INDEX NAME)

RN 841234-61-5 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-[4-[3-(cyclopropylethynyl)phenyl]-3-hydroxybutyl]hydrazino]ethyl]- (9CI) (CA INDEX NAME)

11/564,974

PAGE 1-A

$$C = C \qquad CH_2 - CH_2 -$$

PAGE 1-B

- CO2H

IT 841234-65-9P 841234-67-1P 841234-71-7P

841234-76-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);

RACT

(Reactant or reagent)

(preparation of benzoic acid derivs. having hydrazide moiety as prostaglandin receptors modulators for treatment of asthma, hypertension, etc.)

RN 841234-65-9 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-(3-hydroxybutyl)hydrazino]ethyl]-, methyl

ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{OH} \\ \text{NH-CH}_2-\text{CH}_2-\text{CH-Me} \\ \\ \text{CH}_2-\text{CH}_2-\text{N-Ac} \\ \\ \text{MeO-C} \\ \\ \text{O} \end{array}$$

RN 841234-67-1 CAPLUS

CN Benzoic acid, 4-[2-[1-acetyl-2-(3-hydroxyoctyl)hydrazino]ethyl]-, methyl

ester (9CI) (CA INDEX NAME)

RN 841234-71-7 CAPLUS

CN Benzoic acid,

4-[2-[1-acetyl-2-(3-hydroxyoctyl)-2-methylhydrazino]ethyl]-, methyl ester (9CI) (CA INDEX NAME)

Me OH
$$N-CH_2-CH_2-CH-(CH_2)_4-Me$$
 $N-CH_2-CH_2-CH-(CH_2)_4-Me$ $N-CH_2-CH_2-CH-(CH_2)_4-Me$ $N-CH_2-CH_2-CH-(CH_2)_4-Me$ $N-CH_2-CH_2-CH-(CH_2)_4-Me$

RN 841234-76-2 CAPLUS

CN Benzoic acid,

4-[2-[2-[3-hydroxy-4-[3-(trifluoromethyl)phenyl]butyl]-1-(1-oxopropyl)hydrazino]ethyl]-, methyl ester (9CI) (CA INDEX NAME)

IC ICM C07C243-00

CC 25-17 (Benzene, Its Derivatives, and Condensed Benzenoid Compounds)

Section cross-reference(s): 1, 63

IT 841234-41-1P 841234-42-2P 841234-43-3P

841234-44-4P 841234-45-5P 841234-46-6P

841234-47-7P 841234-49-9P 841234-50-2P

841234-51-3P 841234-52-4P 841234-53-5P

841234-54-6P 841234-55-7P 841234-56-8P

841234-57-9P 841234-58-0P 841234-59-1P

841234-60-4P 841234-61-5P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of benzoic acid derivs. having hydrazide moiety as prostaglandin receptors modulators for treatment of asthma, hypertension, etc.)

IT 136333-97-6P 346672-98-8P 518345-38-5P 841234-62-6P 841234-63-7P

841234-64-8P 841234-65-9P 841234-66-0P 841234-67-1P

841234-68-2P 841234-69-3P 841234-70-6P 841234-71-7P

841234-72-8P 841234-73-9P 841234-74-0P 841234-75-1P

841234-76-2P 841234-77-3P 841234-78-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent)

(preparation of benzoic acid derivs. having hydrazide moiety as prostaglandin receptors modulators for treatment of asthma, hypertension, etc.)

L4 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1979:474573 CAPLUS

DOCUMENT NUMBER:

91:74573

TITLE:

Studies on the syntheses of heterocyclic compounds. 776. Cyclization of N-substituted mandelohydrazide

with formaldehyde

AUTHOR(S):

Kametani, Tetsuji; Kigasawa, Kazuo; Hiiragi,

Mineharu;

Wagatsuma, Nagatoshi; Kohagizawa, Toshitaka; Inoue,

Hitoshi

CORPORATE SOURCE:

Pharm. Inst., Tohoku Univ., Sendai, Japan

SOURCE: -

Yakugaku Zasshi (1979), 99(2), 135-40

CODEN: YKKZAJ; ISSN: 0031-6903

DOCUMENT TYPE:

Journal

LANGUAGE:

Japanese

GI

PhCH(CR¹)CO R N-NR COCH(CR¹)Ph III, R¹=Ac

AB Acid-catalyzed cyclization of N-substituted mandelohydrazides PhCH(OH)CONHNHR (R = Me, cyclopropylmethyl, PhCH2, PhCH2CH2) with paraformaldehyde gave hexahydro-1,2,4,5-tetrazine derivs. I. However, PhCH(OH)CONHNAcCH2Ph (II), when similarly treated, gave

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11/564,974
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cyclization-rearrangement product III. The Ac on the N in II rearranged to alc. O during the cyclization. 68164-66-9P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and rearrangement of) 68164-66-9 CAPLUS RNCN Benzeneacetic acid, α -hydroxy-, 2-acetyl-2-(phenylmethyl)hydrazide (9CI) (CA INDEX NAME) CC 28-22 (Heterocyclic Compounds (More Than One Hetero Atom)) 68164-66-9P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and rearrangement of) ANSWER 4 OF 4 L4CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1978:597482 CAPLUS 89:197482 DOCUMENT NUMBER: TITLE: Cyclization reaction of N-substituted mandelhydrazide with formaldehyde AUTHOR (S₁): Kametani, Tetsuji; Kigasawa, Kazuo; Hiiragi, Mineharu; Wagatsuma, Nagatoshi; Kohagizawa, Toshitaka; Inoue, Hitoshi Pharm. Inst., Tohoku Univ., Sendai, Japan CORPORATE SOURCE: Heterocycles (1978), 9(8), 1031-40 SOURCE: CODEN: HTCYAM; ISSN: 0385-5414 DOCUMENT TYPE: Journal English LANGUAGE.: OTHER SOURCE(S): CASREACT 89:197482 GI ROCHPhCO R1 N COCHPHOR I

AB Tetrazines I (R = H, R1 = Me, cyclopropylmethyl, CH2Ph, CH2CH2Ph; R = Ac,

R1 = CH2Ph) were prepared by cyclocondensation of HOCHPhCONHNHR1 (II) or

HOCHPhCONHNAcCH2Ph (III) with CH2O in the presence of acid. II were prepared by treating HOCHPhCO2Me with R1NHNH2. III was prepared by acetylating II (R1 = CH2Ph) and partial hydrolysis of AcOCHPhCONHNAcCH2Ph.

IT 68164-66-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent)

(preparation and cyclization of, with formaldehyde)

RN 681.64-66-9 CAPLUS

CN Benzeneacetic acid, α -hydroxy-, 2-acetyl-2-(phenylmethyl)hydrazide (9CI) (CA INDEX NAME)

CC 28-22 (Heterocyclic Compounds (More Than One Hetero Atom))
IT 68164-61-4P 68164-62-5P 68164-63-6P 68164-64-7P 68164-66-9P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);

RACT (Reactant or reagent)

(preparation and cyclization of, with formaldehyde)

=> log yCOST IN U.S. DOLLARS SINCE FILE TOTAL SESSION ENTRY 195.22 FULL ESTIMATED COST 22.91 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION -3.12-3.12CA SUBSCRIBER PRICE

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